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(54) ACRYLIC ESTER-BASED RADIATION CURABLE COMPOSITIONS

(71) We, ROHM AND HAAS COMPANY, a corporation organized under the laws of the State of Delaware, United States of America, of Independence Mall West, Philadelphia, Pennsylvania 19105, United States of America, do hereby declare the invention for which we pray that a patent may be granted to us, and the method by which it is to be performed to be particularly described in and by the following statement:—

This invention is concerned with radiation curable compositions comprising isobornyl acrylate and photoinitiator, more particularly radiation curable coating and ink compositions, sheets and the use of isobornyl acrylate as a reactive diluent.

It is well known in the art to deposit polymerizable liquid coatings on substrates and then to dry and cure such coatings by passing them through ovens to crosslink the coatings. It is also known in the art to use coatings which are curable by exposure to actinic radiation such as ultra-violet rays or that from plasma are radiation sources, as well as by exposure to high energy ionizing radiation such as electron beam radiation. These systems have disadvantages in that either ovens are required or that, in radiation curable coatings, oxygen inhibition of the polymerization of the coating may occur, or it may be necessary to employ solvents to decrease the viscosity of the coating for application purposes.

Due to energy and raw material shortages, increasingly strict air pollution standards and safety regulations the search is continuing for one hundred percent polymerizable systems, i.e., compositions which have no highly volatile components but which contain reactive viscosity reducing diluents which become either the sole cured film or part of the cured film. Such types of compositions are known, for example, multifunctional acrylates, methacrylates and itaconates of pentaerythritol, dipentaerythritol and polypentaerythritols and others disclosed in U.S. Pat. Nos. 3,551,235; 3,551,246; 3,551,311; 3,552,986; 3,558,387 and 3,661,614.

Two U.S. patents which disclose radiation cure of monofunctional acrylates are U.S. Pat. No. 3,783,006, which describes a wide number of acrylate monomers for use as diluents in polymer syrups which are cured on metal containers via electron beam radiation and U.S. Pat. No. 3,772,062, which also describes the use of various diluents in the curable coatings.

We have now found that isobornyl acrylate may be used in radiation curable compositions, such as coating compositions, as a reactive diluent in 100% curable systems (i.e. containing only isobornyl acrylate, photoinitiator and optionally at least one other radiation polymerisable component) and may confer on such compositions a desirable balance of properties such as volatility, shrinkage, toxicity and viscosity.

Isobornyl acrylate is a known compound. One method of preparation is disclosed in U.S. Pat. No. 3,087,962.

Low volatility acrylate monomers are available such as trimethylolpropane triacrylate, pentaerythritol triacrylate and neopentylglycol diacrylate; however, these, unlike isobornyl acrylates, cure to highly crosslinked and extremely hard, brittle films. Isobornyl acrylate yields a relatively hard polymer (T_s=94°C.) but unlike the hard multifunctional acrylates does not contribute to increased crosslink density. Higher alkyl acrylates such as lauryl acrylate, isodecyl acrylate and, marginally, 2-ethylhexyl acrylate have the desired lower volatility and viscosity but their use alone as a diluent tends to promote excessive softening. However, in combination with isobornyl acrylate one can achieve an excellent degree of latitude of hardness and flexibility and viscosity control previously unobtainable with other combinations of acrylates. The polymeriza-



5	tion of isobornyl acrylate is accompanied by low volume shrinkage relative to other known reactive diluents thereby aiding and minimizing losses in adhesion and flexibility commonly observed in radiation curable systems. In addition, to these improvements, isobornyl acrylate is of relatively low toxicity, compared to many other ethylenically unsaturated monomers in common use.	5
	According to the invention there is provided radiation curable compositions comprising	
10	 (a) from 1 to 99.9%, preferably from 10 to 98%, by weight of isobornyl acrylate; (b) from 0.1 to 25%, preferably from 1 to 15%, by weight of photoinitiator; (c) from 0 to 80%, by weight of nonradiation polymerisable oligomer, polymer or mixture thereof and, when employed, preferably from 5 to 50% by weight; 	10
	 (d) from 0 to 90% by weight of pigment and, when employed, preferably from 2 to 80% by weight; 	
15	 (e) from 0 to 50% by weight of plasticizer and, when employed, preferably from 5 to 40% by weight; and (f) optionally polymerizable other ethylenically unsaturated monomer, oligomer or polymer or mixture thereof, copolymerizable with isobornyl acrylate and when employed, preferably from 5 to 95% by weight. 	15
20	The ethylenically unsaturated monomers, oligomers and polymers optionally employed with isobornyl acrylate may be any of the known radiation polymerizable coating materials. We have found that isobornyl acrylate is compatible with material such as acrylate pendant polymers, such as acrylated epoxies, acrylated oils, acrylated urethanes, unsaturated polyesters, neopentyl glycol diacrylates; pentaerythritol tri-	20
25	acrylate; trimethylol propane triacrylate; 1,6-hexane diol diacrylate; 2-ethylhexyl acrylate; isodecyl acrylate; hydroxyethyl acrylate; Roskydal 650 polyester (Mobay); Actomer X-70 (an acrylate functional polyester oligomer (Union Carbide)) Epocryl DRH 303.1 and the diacrylate ester of a bisphenol A epoxy resin (Shell). (Roskydal, Mobay, Actomer and Epocryl are Trade Marks).	25
30	In addition to the use of isobornyl acrylate as a reactive diluent at low to modestly high concentration in coating compositions containing reactive resins, crosslinking monomers or other monofunctional monomers, it also has considerable utility as the major if not the sole polymerizable component of a radiation curable coating composition. While very low viscosity may aid in some of these applications, e.g., for ease	30
35	of surface penetration and binding to materials such as wood, leather, paper and cement, it may be desirable to add up to 80% by weight of a non-radiation reactive polymer, oligomer or mixture thereof of certain higher viscosity resins strictly for rheological control, i.e., as "thickeners". The resins do not necessarily have to be reactive in this capacity. Where the hardness of polymerized isobornyl acrylate is not	35
40	required or desired, low viscosity systems of less hardness may be prepared by judicious addition of softer, low viscosity polymers. Examples of these resins include polymers of lower alkyl acrylates and methacrylates wherein the alkyl has from 1—5 carbon atoms such as methyl, ethyl, propyl, butyl and pentyl. If desired, isobornyl acrylate can be made into soft, flexible films by the use of	40
45	inert, nonvolatile plasticizers in the range of from 5 to 50% by weight. Plasticizers which can be employed include phthalate esters such as dibutyl, dioctyl, di-2-ethylhexyl, butylbenzyl diisodecyl esters, Santicizer 262 (Monsanto) and adipate esters such as dioctyl (Santicizer is a Trade Mark); phosphates such as tributoxyethyl, tributyl and tricresyl; phthalyl glycolates such as butyl phthalyl butyl glycolate; and	45
50	sulfonamides such as N-ethyl-o-p-toluenesulfamide; also included are castor oil and its derivatives and polymeric types such as those disclosed in the Rohm and Haas Company booklet "Paraplex and Monoplex Plasticizers—summary of physical performance properties" published May 1968. (Paraplex and Monoplex are Trade Marks). The acrylate based photocurable monomers, polymers and oligomers are generally	50
55	preferred because of higher cure speed. The ethylenically unsaturated portion of the composition other than isobornyl acrylate may comprise 0—98.9% by weight of the composition of one or more acrylate monomers or oligomers or polymers thereof; preferably at least one monomer (or an oligomer or polymer derived from said monomer) of the formula:	55

 $H_2C = C - C - C - R^1$ (I) 60

wherein R is hydrogen or methyl and R' is (C_1-C_{1a}) alkyl preferably (C_1-C_{1a}) alkyl, such as methyl, ethyl, propyl, isopropyl, butyl, hexyl, 2-ethylhexyl, decyl, isodecyl, dodecyl and octadecyl, aryl, for example, mononuclear aryl, such as benzyl and phenyl, cycloalkyl, for example, cycloalkyl of from 5 to 12 nuclear carbon atoms such as cyclohexyl, trimethyl cyclohexyl, cyclopentyl, cycloctyl and dicyclopentenyl, substituted aryl or substituted cycloalkyl wherein the substituent is alkyl, halo and/or hydroxy, or \mathbb{R}^1 is

$$-\left(R^2-0\right)_zR^3$$
 (II)

wherein R² is an alkylene radical of from 1 to 8 carbon atoms such as methylene, ethylene, propylene or isobutylene; z is a whole number from 2 to 25 and R³ is as defined above for R¹, or R¹ is:

$$\frac{\left(R^2 - co - o \right)_2}{2} R^3 \tag{III}$$

wherein R2, R2 and z are as described above.

Typical of the acrylic monomers which may be employed in this invention include monoacrylates, such as methyl acrylate, isopropyl acrylate, cyclopentyl acrylate, 2-ethylhexyl acrylate, decyl acrylate, isodecyl acrylate, decyl thioacrylate, dodecyl acrylate, octadecyl acrylate, hydroxyethyl acrylate and hydroxybutyl acrylate; diacrylates, such as ethylene glycol diacrylate, triethylene glycol diacrylate, tetraethylene glycol diacrylate, neopentyl glycol diacrylate, 1,6-hexane diol diacrylate, polypropylene glycol diacrylate, 1,3-butane diol diacrylate and 1,4-butane diol diacrylate; triacrylates such as trimethylol propane triacrylate and pentaerythritol triacrylate or tetraacrylates such as pentaerythritol tetraacrylate.

A photoinitiator or sensitizer is added to the composition before exposing the composition to radiation. The photoinitiators or sensitizers are employed in the amounts of from 0.1 to 25% by weight of the composition and preferably from 1 to 15% by weight of the total polymerizable composition. Preferred photoinitiators employed include selected acyloins or derivatives thereof, for example, benzoin alkyl ethers such as benzoin methyl ether and benzoin ethyl ether, desyl halides such as desyl bromide and desyl chloride, desyl amine, benzophenone derivatives, acetophenone compounds, polychlorinated aromatic compounds, a combination of organic carbonyls and amines or mixtures thereof. The acetophenone photoinitiators are disclosed in U.S. Pat. No. 3,715,293. The combination of organic carbonyls and amines is disclosed in U.S. Pat. No. 3,795,807.

The invention also provides a method of coating which comprises applying a composition of the invention to a substrate and curing or drying the coating by exposure to radiation.

The compositions of this invention are dried or cured by exposure to radiation. The compositions exhibit maximum sensitivity in the range of from about 1800 to 5000 A and any source of actinic light can be employed. Suitable sources of radiation include electron beams, carbon arcs, mercury-vapor arcs, fluorescent lamps with ultraviolet light emitting phosphors, argon glow lamps, photographic flood lamps, Van der Graaf accelerators, resonant transformers, betatron linear accelerators, gamma radiation emitters and combinations thereof.

The photopolymerizable coating compositions of the present invention are useful as coatings such as adhesives, markers, vehicles for printing inks, lacquers and paints on various substrates including metals ceramic, cement, wood, plastics, textiles, paper, floor tiles, glass, roads, parking lots and airfields. The compositions are also useful in the preparation of photopolymerizable elements, i.e., a support having disposed thereon a photopolymerizable layer of a composition as described herein. Moreover, various dyestuffs, pigments, plasticizers, lubricants and other modifiers may be incorporated to obtain certain desired characteristics in the finished products.

When a photopolymerizable composition of the present invention is used as an adhesive, at least one of the lamina must be translucent when ultra-violet light is used. When the radiation source is an electron beam or gamma radiation at least one of the lamina must be capable of transmitting high energy electrons or gamma radiation respectively, and neither is necessarily translucent to light. Typical laminations include polymer coated Cellophane to polymer coated Cellophane films, treated poly-

4	1,550,382	4
	ethylene to treated polyethylene films, Mylar polyester film to a metal substrate such as copper and the like, opaque oriented polypropylene to aluminum or polymer coated Cellophane to polypropylene. (Mylar is a Registered Trade Mark).	
5	metal coatings such as in coil coating and in metal decorating. For example, in metal decorating isobornyl acrylate is useful in formulating pigmented base coats, inks and clear overprint varnishes. Glass and plastics may also be coated and the coatings are conventionally applied by dip coating direct and reverse and processes.	5
10	Cellophane; glass, treated and untreated polyethylene, for example, in the form of disposable cups or bottles. (Cellophane is a Registered Trade Mark). Examples of metals which may be coated include sized and unsized tin plate, tin free steel and aluminum.	iO
15	The compositions may be pigmented with organic or inorganic pigments, for example, molybdate orange, titanium white, chrome yellow, phthalocyanine blue and carbon black, as well as colored with dyes. Stock which may be printed includes paper, clay coated paper and boxboard. In addition, the compositions of the present invention are suitable for the treatment of textiles, both natural and synthetic, for example, in yehicles for textile printing into an formatical linear and synthetic, for example, in	15
20	vehicles for textile printing inks or for specialized treatments of fabrics to produce water repellency, oil and stain resistance, crease resistance and the like. A general ink formulation would be the vehicle of the composition of 1—99.8% by weight of isobornyl acrylate, 0—98.8% by weight of an ethylenically unsaturated monomer, oligomer or polymer or mixture thereof, 0.1—25% by weight of photoinitiator and 0.1—90% by weight of pigment based on the total vehicle ink composition.	20
25	a sheet or plate, having superimposed thereon a layer of the above described photo- polymerizable compositions. Suitable base or support materials include metals, for example, steel and aluminum plates, sheets and foils. Also films or plates composed of various film forming synthetic resins or high polymers, such as addition polymers, and	25
30	polymers; vinylidene chloride copolymers with either vinyl chloride, vinyl acetate or acrylonitrile; and vinyl chloride copolymers with vinyl acetate or acrylonitrile; linear condensation polymers such as polyesters, for example, polyethylene, teraphylenes	30
35	and polyamides. Fillers or reinforcing agents can be present in the synthetic resin or polymer bases. In addition, highly reflective bases may be treated to absorb ultraviolet light or a light absorptive layer can be transposed between the base and photopolymerizable layer. Photopolymerizable elements can be made by exposing to ultra-violet light selected portions of the absorptive layer.	35
40	selected portions of the photopolymerizable layer thereof until addition polymerization is completed to the desired depth in the exposed portions. The unexposed portions of the layer are then removed, for example, by use of solvents which dissolve the monomer or prepolymer but not the polymer. As stated above, any suitable source of actinic energy may be used, for example a 200 wart lineal inch medium are suitable source.	40
45	which a photopolymerizable composition will dry include the specific ingredients in the composition, concentration of the photoinitiators, thickness of the material, nature and intensity of the radiation source and its distance from the material, the presence or absence of oxygen and the ambient temperature.	45
50	The compositions of the present invention are generally employed as films having a thickness of from 0.1 to 30 mils, and preferably from 0.2 to 10 mils. One embodiment of the invention provides sheets prepared from the radiation curable compositions. The sheets may have a thickness of from 30 to 1000 mils. Some preferred embodiments of the invention will now be more particularly	50
	weight unless otherwise stated. Example 8 is a comparative Example. The following is a list of either how various tests were conducted or a reference to where the tests are described:	55
60	Viscosity— Where reported in centipoise (cps.), viscosities were determined with a Brookfield Viscometer at room temperature. "Paint Testing Manual", Gardener and Sword, 12th edition, 1962, p. 178. Where reported in centistokes, viscosities were determined with Gardner Bubble Standards. "Paint Testing Manual", Gardener and Sword, 12th edition, 1962, p. 172.	60

Penmerythritol triacrylate

Methyl diethanolamine

Benzophenone

55

20

2

55

	-			,,				6
. 5	The liquid co panels by a wire we the procedure described excellent mar resist 150°F, for one hou	ribed in]	Evample 1	The suck	iess of U.Z)	mils and t	hen cured by	5
			FYA	MDIES				ر
10	EXAMPLE 3 Coating of Isobornyl acrylate and Epocryl DRH-303.1 A 100% polymerizable liquid coating is prepared by mixing until clear and homogeneous Epocryl DRH-303.1, a high viscosity diacrylate ester of a bisphenol A epoxy resin (Shell), with isobornyl acrylate and diethoxyacetophenone (photosensitizer) in the proportions given below.							
	Coating C	ompositio	n					
	Epocryl D				Part	s by Wei	ght	
	Isobornyl a	1011-303. <u>[</u> CTV 210			•	45		
	Diethoxyac	etophenor				55		
	•	_				2		
15	The liquid coar aluminum test panels cured by the proced 2H) has very good n at 150°F, for 1 hou	ure descri	ihed in Ev	to yield a I	nm mickness	ot 0.25 m	uls and then	15
20			EVA	MDIE				
	· Com	parison o	of Isoborny	MPLE 4	with Styrene			20
			Isobornyi	methacryli	200			
25	Isobornyl methacrylate The utility of isobornyl acrylate as a diluent monomer for 100% polymerizable unsaturated polyester coatings conventionally employing styrene diluent is afforded by the comparative cure results of Table II. Therein, the unsaturated polyester resin Roskydal 650 (Mobay) was dissolved in styrene, isobornyl acrylate, as well as in isobornyl methacrylate to demonstrate the cure speed advantage of the preferred acrylate moiety. The coatings were cast to yield 1.5 mil films on aluminum test panels and cured as described in Example 1.							25
30			ample 1.					
30	_		TAE	BLE II	•			
	Coating Composition						•	30
	(parts by weight)	A	В	С	D	E	F	
	Roskydal-650	0	0	0	65	65		
26	Styrene	100	Ö	ŏ	35	0	65 0	
35	Isobornyl acrylate	0	100	Ŏ	ō	35	ŏ	2.5
•	Isobornyl			•	J	٠.	U	35
	methacrylate	0	0	100	0	0	35	
	Benzoinethyl ether	3	3	3	3	. 3	3	
40	Cured Film Properties					•		
	Appearance	evaporate	es cures	no cure	some cure	cures	some cure	40
	Pencil Hardness	1		liquid	heavy mar	mar free	heavy mar	
•	_	NT*	3 H	NT	NT	F	NT	
	* not tested							
			EXAM	IPLE 5				
45	Cor	noosition	of Isoborn	vi acrylate	, 2-Ethylhex	-vl		4.5
	•	acı	viate and	Actomer X	C-80			45
	A 100% polymer until clear and homog	izable coa	ting is pre	pared by a	dmixing the	following:	ingredients tes:	
	Coating Con				_			
50	Actomer X-8	-			1 at the	by Weigh	ıL	
	Isobornyl ac					60		50
	2-Ethylhexyl	acrylate				20		
	Methyl dieth	anolamina				20		
	Benzophenon	C	•			3 2		
6.5				_		_		
55	The liquid coating mercial vinyl asbestos a glossy, resilient, mar	TYPE LOOF	ing tile and	cured as a	described in	Evample 1	e of com- l to afford	55

	EXAMPLE	6			•			
	Isobornyl acrylate with Nonradia	tion Rea	ctive	Polyr	ners			
	The compositions of Table III are prepare	d by adm	nixing	the c	'AMBABE	nts in c	OD-	
	vermonar equipment until clear and nomogeneo	us. Ine	wet co	natino	C 278 27	mlied u	rith	
	a No. 4 wire wound rod to aluminum test pane	ls (0.25	mil fil	m rh	ickness)	and on	rad	
5	as described in Example 1.	(0.22			icances)	and cu.	i Cu	
								5
	TABLE II	T						
	Coating Composition .	•						
	(parts by weight)		Α	В	С	-		
					_	D		
	Isobornyl acrylate	1	100	20	20	20		
	Isobutyl methacrylate polymer							
10	(Mw 80,000; Tg 50°C.)		•		_	_		
10	Methyl methografica 70 (Erbyl comics 30 co		0	80	0	0		10
	Methyl methacrylate 70/Ethyl acrylate 30 co	polymer-		•				
	$(\overline{M}w=90,000; Tg 60^{\circ}C.)$		_		••			
	Mathyl mathematics 55 (Ethal and and a	. •	0	0	80	0		
	Methyl methacrylate 55/Ethyl acrylate 45 co	polymer-						
	$(\overline{M}w = 70,000; Tg 35^{\circ}C.)$			^	•			
15	Benzoinethyl ether		0 3	0	0	80		
	System Properties)	3	3	3		. 15
	Liquid viscosity (centipoise)		7	240	2700	0.00		
	Cured Film Pencil Hardness		7 3 H	240 H	3700	960		
	Carca 1 mm 1 cach 11atmess)II	п	2 H	H		
	EXAMPLE	7						
20	Coating of Concrete		.d					
	Composition A of Example 6 is applied by	wining a	ou ohin 1	Inna-				20
	slab and a wood test slab. The slabs were then	wibnig g	- eke	MACE	OH # CO	icrete to	=st	
	in Example 1 to yield water repellent, protectiv	Lured 6	y we	broce	cuire as	describ	ea	
	in parapre 1 to large again telegram brothers	e nara m	тэпез	OH D	orr and	strates.		
	EVANDE	0						
25	EXAMPLE Page of Vilority							0.2
23	Rates of Volatili	zauon						25
	The volatility characteristics of isobornyl ac	Tylate 15	comp	area '	WILL OLD	er anne	nt	
	monomers. Seven (7) cm. diameter circles of W	naturan r	40. L	micr	paper is	saturat	ea	
•	with monomer (ca. 1—1.3 gm.) and weight loss is specimens being placed in a laboratory hood of		red as	uri B	CHOH OF	ume, t	DC.	
30	Registered Trade Mark). Weight loss versus time	in linear	air v	CIUCIL	у. (WДа	Of less	; a	20
50	the samples. The resulting rates of weight loss,	is inter	MIOU.	gu at millio	15451 00	% 10SS	OL	30
	from the specified samples are given below.	expresse	и ш	mmi	rams be	t tumnin	.c3	
	from the specifical samples are given below.							
	• •			Rote /	of Volati	lization		
	Monomer				ng./min			
				(4		•)		
35	Styrene				19	• •		35
	Butyl acrylate				17			
	Cyclohexyl acrylate				1.9			
	2-Ethylhexyl acrylate	•			0.5			
	Isobornyi acrylate				0.25			40
40	Isodecył acrylate				0.1			40
	EXAMPLE	9 .						
	Isobornyl acrylate Composition		Plastic	izers				
	Composition							
	(parts by weight)	Α		В		С		
45	Isobomyi acrylate	100		95		90		45
73	Santicizer 262 ¹	0		5		10		7.7
	2,2-Diethoxyacetophenone	2		2		2		
	Cured Film Properties	L		-		-		
	Knoop Hardness	14		10		4.4		
50	Mandrel Flexibility (diameter passed, in.)	>6		>6		4		50
J U	•					•		50
	¹ Santicizer 262 is a monomeric phthalate type	plasticize	r (Mo	nsant	o) of ur	adisclose	ed ·	
	composition having a molecular weight of 3	96.5	•		-			

	WHAT WE CLAIM IS:—	
	1. A radiation curable composition comprising	
	(a) from 1 to 99.9% by weight of isobornyl acrylate;	
	(b) from 0.1 to 25% by weight of a photoinitiator;	
5	(c) 0 to 80% by weight of nonradiation polymerizable oligomer, polymer or	5
•	mixture thereof;	
	(d) from 0 to 90% by weight of pigment;	
	(e) from 0 to 50% by weight of plasticizer; and	
10		10
10	oligomer or polymer or mixture thereof copolymerizable with isobornyl	10
	actylate.	
	2. A composition as claimed in Claim 1 comprising:	
	(a) from 10 to 98% by weight of isobornyl acrylate; and	
15	(b) from 1 to 15% by weight of photoinitiator.	15
13	3. A composition as claimed in claim 1 or 2 which is in the form of a 100% radiation polymerizable composition which contains only components (a) and (b) and	ر ،
	optionally component (f).	
	4. A composition as claimed in any preceding claim wherein the photoinitiator comprises an acyloin or derivative thereof; desyl halide, desyl amine, benzophenone	
20		20
20	derivative, acetophenone compound, polychlorinated compound, a combination of	20
	organic carbonyl and amine or mixture thereof.	
	5. A composition as claimed in any preceding claim containing from 5 to 95% by weight of radiation polymerizable other ethylenically unsaturated monomer, oligomer	
25	or polymer or mixture thereof copolymerizable with isobornyl acrylate. 6. A composition as claimed in any of Claims 1, 2 or 4 containing from 5 to 50%	25
23	by weight of nonradiation polymerizable oligomer, polymer or mixture thereof.	23
	7. A composition as claimed in any of Claims 1, 2 or 4 containing from 2 to	
	80% by weight of pigment.	
	8. A composition as claimed in any of Claims 1, 2 or 4 containing from 5 to 40%	
30		30
30	by weight of plasticizer. 9. A composition as claimed in Claim 1 substantially as described in any of the	30
	foregoing Examples 1—7 and 9.	
	10. A method of coating which comprises applying a composition according to	
	any of Claims 1 to 9 to a substrate and curing or drying the coating by exposure to	
35	radiation.	35
33	11. A method as claimed in Claim 10 wherein the coating is dried or cured by	,,,
	electron beam.	
	12. A method as claimed in Claim 10 wherein the coating is dried or cured by	
	actinic light.	
40	13. A substrate having thereon a radiation cured composition according to any of	40
+∪	Claims 1—9.	
	14. A substrate as claimed in Claim 13 wherein the cured composition is from 0.1	
	to 30 mils thick.	
	15. A substrate as claimed in Claim 13 wherein the cured composition is from 0.2	
45	to 10 mils thick.	45
TJ	16. A sheet prepared from a radiation cured composition according to any of	
	Claims 1—9.	
	17 A sheet as claimed in Claim 16 having a thickness of 30 to 1000 mils.	

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